

Substitute for form 1449A/PTO

INFORMATION DISCLOSURE
STATEMENT BY APPLICANT

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Sheet 1 of 3

Complete if Known

Application Number	10/806,296
Filing Date	March 22, 2004
First Named Inventor	Rabi, J.A.
Group Art Unit	1623
Examiner Name	Unassigned
Attorney Docket Number	06171.105107 IDX 1012B US

3478701_2.DOC

U.S. PATENT DOCUMENTS

Examiner Initials *	Cite No. ¹	U.S. Patent Document		Name of Patentee or Applicant of Cited Document	Date of Publication of Cited Document MM-DD-YYYY	Pages, Columns, Lines, Where Relevant Passages/Relevant Figures Appear
		Number	Kind Code (if known)			
AA		3,891,623	A	Verbruggen et al.	06-24-1975	
AB		4,754,026	A	Kawada et al.	06-28-1988	
AC		4,957,924	A	Beauchamp	09-18-1990	
AD		5,144,018	A	Kuzuhara et al.	09-01-1992	all
AE		5,565,438	A	Chu et al.	10-15-1996	
AF		5,567,688	A	Chu et al.	10-23-1996	duplicates
AG		5,587,362	A	Chu et al.	12-24-1996	
AH		6,153,594	A	Børretzen et al.	11-28-2000	
AI		6,248,878	B1	Matulic-Adamic et al.	06-19-2001	
AJ		6,271,212	B1	Chu et al.	08-07-2001	
AK		6,395,716	B1	Gosselin et al.	05-28-2002	
AL		6,444,652	B1	Gosselin et al.	09-03-2002	
AM		2003-0050229	A1	Sommadossi et al.	03-13-2003	
AN		6,566,344	B1	Gosselin et al.	05-20-2003	
AO		6,569,837	B1	Gosselin et al.	05-27-2003	
AP		2003-0083306	A1	Imbach et al.	05-01-2003	
AQ		2004-0006002	A1	Sommadossi et al.	01-08-2004	

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		Office ³	Number	Kind Code ² (if known)			
AR		DD	0,140,254	Z	Akad. Wissenschaft der DDR Zentralinstitut für Molekularbiologie	02-20-1980	
AS		DE	42 24 737	A1	Prof. Dr. Herbert Schott	02-03-1994	all
AT		EP	0 352 248	A1	Johansson et al.	01-24-1990	
AU		JP	62-93645	A	Jpn. Kokai Tokkyo Koho	10-21-1994	duplicates translation
AV		WO	95/07287	A1	CNRS	03-16-1995	
AW		WO	96/11204	A1	Max Delbrück Centr. Mol. Med.	04-18-1996	
AX		WO	96/13512	A2	Genencor Int'l; Lipitek	05-09-1996	
AY		WO	96/40164	A1	Emory U.; UAB Res. Found.; CNRS	12-19-1996	
AZ		WO	00/09531	A2	Novirio Pharm. [Idenix]; CNRS	02-24-2000	
AAA		WO	01/90121	A2	Novirio [Idenix]; Univ.... Cagliari	11-29-2000	
AAB		WO	01/96353	A2	Novirio Pharm. [Idenix]; C.N.R.S.	21-20-2001	

Examiner Signature	G. Karishma	Date Considered	9/29/06
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OTHER PRIOR ART - NON PATENT LITERATURE DOCUMENTS

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BA	BA	BENZARIA, S., et al., "Synthesis of potential prodrugs of b-4-L-dC, a potent and selective anti-HBV agent," <i>Antiviral Res.</i> , 50:A79 (2001). [Abstract no. 137].	
BB	BB	BLOCH, A., et al. "The Role Of The 5'-Hydroxyl Group Of Adenosine In Determining Substrate Specificity For Adenosine Deaminase," <i>J. Med. Chem.</i> , 10(5):908-12 (September 1967).	
BC	BC	BRYANT, M.L., et al., "Antiviral L-nucleosides specific for hepatitis B virus infection," <i>Antimicrob. Agents Chemother.</i> , 45(1):229-235 (January 2001).	
BD	BD	BUDAVARI, et al., <i>The Merck Index</i> , 12th Edition, Entry no. 10039, p. 10044.	
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BF	BF	CRETTON-SCOTT, E., et al., "Pharmacokinetics of β-L-2'-deoxycytidine prodrugs in monkeys," <i>Antiviral Res.</i> , 50:A44 (2001) [Abstract no. 16].	
BG	BG	DAVISON, V.J., et al., "Synthesis of Nucleotide 5'-Diphosphates from 5'-O-Tosyl Nucleosides," <i>J. Org. Chem.</i> , 52(9):1794-1801 (1987).	
BH	BH	FOX, J.J., et al., "Thiolation of nucleosides. II. Synthesis of 5-methyl-2'-deoxycytidine and related pyrimidine nucleosides," <i>J. Am. Chem. Soc.</i> , 81:178-187 (1959).	
BI	BI	HOARD, D.E., et al., "Conversion of Mono- and Oligodeoxyribonucleotides to 5'-Triphosphates," <i>J. Am. Chem. Soc.</i> , 87(8):1785-1788 (April 20, 1965).	
BJ	BJ	HOLY, A., "Nucleic Acid Components and Their Analogs. CLIII. Preparation of 2'-deoxy-L-Ribonucleosides of the Pyrimidine Series," <i>Collect. Czech. Chem. Commun.</i> , 37(12):4072-4087 (1972).	
BK	BK	IMAI, K., et al., "Studies on Phosphorylation. IV. Selective Phosphorylation of the Primary Hydroxyl Group in Nucleosides." <i>J. Org. Chem.</i> , 34(6):1547-1550 (June 1969).	
BL	BL	KANEKO, M., et al., "A convenient synthesis of cytosine nucleosides," <i>Chem. Pharm. Bull.</i> , 20:1050-1053 (1972).	
BM	BM	KERR, S.G., et al., "N4-(dialkylamino)methylene derivatives of 2'-deoxycytidine and arabinocytidine: physicochemical studies for potential prodrug applications," <i>J. Pharm. Sci.</i> , 83(4):582-586 (April 1994).	
BN	BN	LIN, T.-S., et al., "Synthesis of Several Pyrimidine L-Nucleoside Analogues as Potential Antiviral Agents," <i>Tetrahedron Letters</i> , 51(4):1055-1068 (1995).	
BO	BO	LUH, T.-Y., et al., "A convenient method for the selective esterification of amino-alcohols," <i>Synthetic Communications</i> , 8(5):327-333 (1978).	
BP	BP	MAGA, Giovanni, et al., "Lack of stereospecificity of suis pseudorabies virus thymidine kinase," <i>Biochem. J.</i> , 294(2):381-385 (1993).	
BQ	BQ	McCORMICK, J., et al., "Structure and total synthesis of HF-7, a neuroactive glyconucleoside disulfate from the funnel-web spider <i>Hololenia curta</i> ," <i>J. Am. Chem. Soc.</i> , 121(24):5661-5664 (1999).	

all duplicates

Examiner Signature	G. Krizil	Date Considered	7/29/06
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Attorney Docket Number	06171.105107 IDX 1012B US

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CB	STANDRING, D.N., et al., "Antiviral beta-L-nucleosides specific for hepatitis B virus infection," <i>Antiviral Chem. & Chemother.</i> , 12 (Suppl. 1):119-129 (2001).		
CC	TANG, X.-Q., et al., "2'-C-Branched ribonucleosides: Synthesis of the phosphoramidite derivatives of 2'-C-b-methylcytidine and their incorporation into oligonucleotides," <i>J. Org. Chem.</i> , 64(3):747-754 (1999).		
CD	TYRSTED, G., et al. "Inhibition of the synthesis of 5-phosphoribosyl-1-pyrophosphate by 3'-deoxyadenosine and structurally related nucleoside analogs." <i>Biochim. Biophys. Acta</i> , 155(2):619-622 (February 26, 1968).		
CE	VERRI, A., et al., "Lack of enantiospecificity of human 2'-deoxycytidine kinase: relevance for the activation of β-L-deoxycytidine analogs as antineoplastic and antiviral agents," <i>Molecular Pharmacology</i> , 51(1):132-138 (January 1997).		
CF	VERRI, A., et al., "Relaxed Enantioselectivity of Human Mitochondrial Thymidine Kinase and Chemo-therapeutic Uses of L-Nucleoside Analogues," <i>Biochem. J.</i> , 328(1):317-320 (November 15, 1997).		
CG	Von JANTA-LIPINSKI, M., et al., "Newly Synthesized L-Enantiomers of 3'-Fluoro-Modified β-2'-Deoxyribonucleoside 5'-Triphosphates Inhibit Hepatitis B DNA Polymerase but not the Five Cellular DNA Polymerases α, β, γ, δ, and ε Nor HIV-1 Reverse Transcriptase," <i>J. Medicinal Chemistry</i> , 41(12):2040-2046 (May 21, 1998).		
CH	ZEDECK, M.S., et al., "Inhibition of the steroid-induced synthesis of D5-3-ketosteroid isomerase in <i>Pseudomonas testosterone</i> by a new purine deoxyribonucleoside analog: 6-chloro-8-aza-9-cyclopentylpurine," <i>Mol. Pharmacol.</i> , 3(4):386-395 (1967).		
CI	ZHANG, W., et al., "Removal of silyl protecting groups from hydroxyl functions with ammonium fluoride in methanol," <i>Tetrahedron Letters</i> , 33:1177-1180 (1992).		

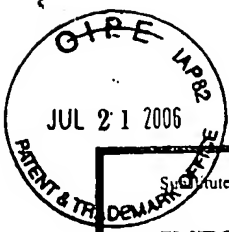
all duplicates -

Examiner Signature	G. Kizilhan	Date Considered	9/29/06
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Sheet

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Application Number	10/806,296
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First Named Inventor	Rabi, J.A.
Group Art Unit	1623
Examiner Name	Ganapathy Krishnan

Attorney Docket Number

06171.105107 IDX 1012C

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U.S. PATENT DOCUMENTS

Examiner Initials *	Cite No. ¹	U.S. Patent Document		Name of Patentee or Applicant of Cited Document	Date of Publication of Cited Document MM-DD-YYYY	Pages, Columns, Lines, Where Relevant Passages/Relevant Figures Appear
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CK	AA	3,891,623	A	Vorbruggen et al.	06-24-1975	
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	AC	4,957,924	A	Beauchamp	09-18-1990	
	AD	5,144,018	A	Kuzuhara et al.	09-01-1992	
	AE	5,565,438	A	Chu et al.	10-15-1996	
	AF	5,567,688	A	Chu et al.	10-23-1996	
	AG	5,587,362	A	Chu et al.	12-24-1996	
	AH	6,153,594	A	Børretzen et al.	11-28-2000	
	AI	6,248,878	B1	Matulic-Adamic et al.	06-19-2001	
	AJ	6,271,212	B1	Chu et al.	08-07-2001	
	AK	6,395,716	B1	Gosselin et al.	05-28-2002	
	AL	6,444,652	B1	Gosselin et al.	09-03-2002	
	AM	2003-0050229	A1	Sommadossi et al.	03-13-2003	
	AN	6,566,344	B1	Gosselin et al.	05-20-2003	
	AO	6,569,837	B1	Gosselin et al.	05-27-2003	
	AP	2003-0083306	A1	Imbach et al.	05-01-2003	
	AQ	2004-0006002	A1	Sommadossi et al.	01-08-2004	
CK		2005/0020825		Storer et al.	01-27-2005	

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		Office ³	Number	Kind Code ² (if known)				
CK	AR	DD	0,140,254	Z	Akad. Wissenschaft der DDR Zentralinstitut für Molekularbiologie	02-20-1980		
	AS	DE	42 24 737	A1	Prof. Dr. Herbert Schott	02-03-1994	with English abstract	
	AT	EP	0 352 248	A1	Johansson et al.	01-24-1990		
	AU	JP	62-93645	A	Jpn. Kokai Tokkyo Koho	10-21-1994	machine translation	
	AV	JP	07224081	A2	Kobayashi Perfumery Co.	08-22-1995	English translation	
	AW	JP	2000290289	A2	Mitsui Chemicals Inc.	10-17-2000	English translation	
	AX	WO	95/07287	A1	CNRS	03-16-1995		
	AY	WO	96/11204	A1	Max Delbrück Centr. Mol. Med.	04-18-1996	with English abstract	
	AZ	WO	96/13512	A2	Genencor Int'l; Lipitek	05-09-1996		
CK	AAA	WO	96/40164	A1	Emory U.; UAB Res. Found.; CNRS	12-19-1996		

Examiner Signature	Gr. Krishnan	Date Considered	9/29/06
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	BL	HOARD, D.E., et al., "Conversion of Mono- and Oligodeoxyribonucleotides to 5'-Triphosphates," <i>J. Am. Chem. Soc.</i> , 87(8):1785-1788 (April 20, 1965).	
	BM	HOLY, A., "Nucleic Acid Components and Their Analogs. CLIII. Preparation of 2'-deoxy-L-Ribonucleosides of the Pyrimidine Series," <i>Collect. Czech. Chem. Commun.</i> , 37(12):4072-4087 (1972).	
	BN	HUBBARD, A.J. et al., "An Investigation by 1H NMR Spectroscopy Into the Factors Determining the β : α Ratio of the Product in 2'-Deoxynucleoside Sythesis", <i>Nucleic Acids Research</i> , 12(7) : 6827-6837(1984)	
	BO	IMAI, K., et al., "Studies on Phosphorylation. IV. Selective Phosphorylation of the Primary Hydroxyl Group in Nucleosides." <i>J. Org. Chem.</i> , 34(6):1547-1550 (June 1969).	
CK	BP	KANEKO, M., et al., "A convenient synthesis of cytosine nucleosides," <i>Chem. Pharm. Bull.</i> , 20:1050-1053 (1972).	

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	CG	STANDRING, D.N., et al., "Antiviral beta-L-nucleosides specific for hepatitis B virus infection," <i>Antiviral Chem. & Chemother.</i> , 12 (Suppl. 1):119-129 (2001).	
	CH	TANG, X.-Q., et al., "2'-C-Branched ribonucleosides: Synthesis of the phosphoramidite derivatives of 2'-C-b-methylcytidine and their incorporation into oligonucleotides," <i>J. Org. Chem.</i> , 64(3):747-754 (1999).	
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